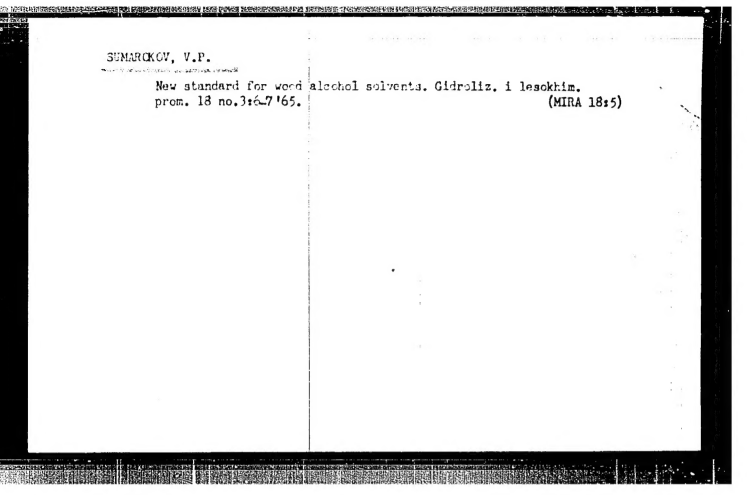


Production of high prom. 17 no.7:16-1	-quality methylacetate. G	idroliz. i lesokhim. (MIRA 17:11)	
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SUMAROKOV, Viktor Favlovich
Anna 11'inichna; TULL'AKOV, B.V., red.

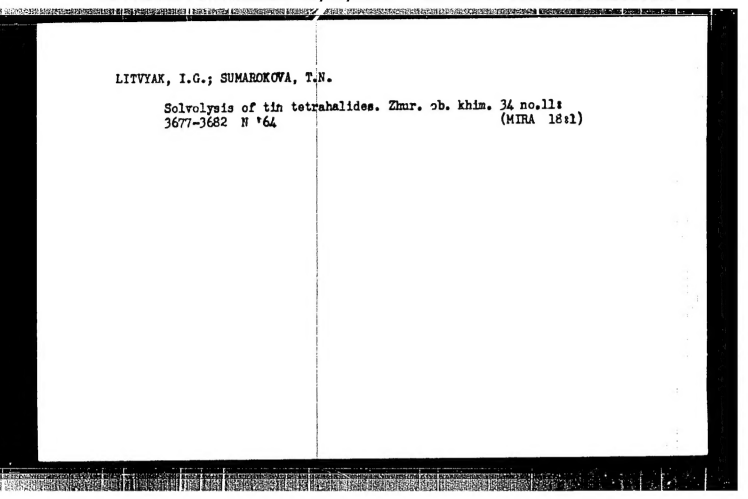
[Tall oil] Tallovoe maslo. Moskva, Lesnaia promyshlennost', (MIRA 18:3)

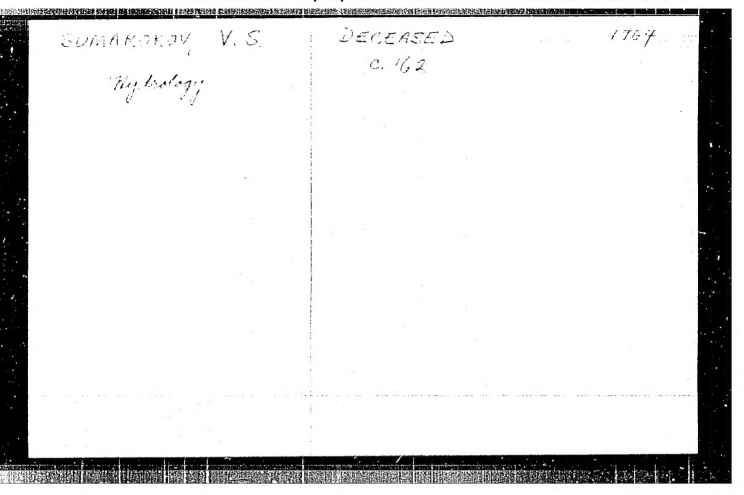


RONDRAT' YEVA, Ye.N.; TARANENKO, L.I.; SUMARUKOVA, R.S.

Requirement of some microelements by purple and green sulfur bacteria. Nauch. dokl. vys. shkoly; biol. nauki no.2:176-180 (MIRA 18:5)

1. Rekomendovana kafedroy mikrobiologii Moskovskogo gousdarstvennogo universiteta im. M.V. Lomonosova.





Binary Systems Formed by SnCll, SbCl3, and AsCl3. The System Stoll-Col1900H, T. Shumarokova, i. Usanovich, Lab Phys Chem, Inst Chem, Acad Sci Azakh SSR Zhur Obshch Khim" Vol XXI, No 7, 1219-1222 Zhur Obshch Khim" Vol XXI, No 7, 1219-1222 Zhur Obshch Khim Vol XXI, No 7, 1219-1222 This cosity-diagrams showed presence of scid-base interaction between components and of compds SbCl3 - CCl3COOH. SbCl3 - CCl_COOH and 2SbCl3 - CCl3COOH. Dystectic max on fusibility diagram of system (Contd) Tepresented compd SbCl3 - CCl3COOH (mp 56°). This is lst established case of CCl3COOH acting as additive or oxonium base. 191718	SIMAROKOVA, K.	STOCKER A ASSESSMENT STOCKER TO SEE	和中国的国际公司 [2] [2] [2] [3] [3] [4] [4] [4] [4] [4] [4] [4] [4] [4] [4		191718
	grater	. 100	viscosity, density of sy 50, 60, 70°C. Elec cond showed presence of acid-n components and of compond 25°Cl3 · CCl3COOH.	"Binary Systems Formed by SnCl ₄ , SbCl ₃ , and AsCl ₃ , V. The System SbCl ₃ -CCl ₃ COOH;" T. Sumarokova, M. Usanovich, Lab Phys Chem, Inst Chem, Acad Sci Kazakh SSR "Zhur Obshch Khim" Vol XXI, No 7, 1219-1222	

SUMAROKOVA, M.Ya., dotsent.

Apt.delo 2 no.5:51-52 S-0 '53.

Work of the scientific student circle attached to the chair of pharmacognosy of the Chervenkov Academy of Medicine in the academic year 1951-52 (Bulgaria).

(MLRA 6:10)

(Bulgaria--Pharmacognosy--Study and teaching)
(Study and teaching--Pharmacognosy--Bulgaria)

SUMAROKOVA, M.Ya., dotsent; OAVRILTUK, Ys.Ya.

From the pages of the foreign pharmaceutical press (Csechoslovakia). Apt. delo no.4:74-79 JI-Ag '53. (MERA 6:8) (Sechoslovakia--Pharmacy) (Pharmacy--Bibliography) (Bibliography--Pharmacy)

SUMAROKOVA, M.Ya., dotsent

Pifth edition of the Hungarian Pharmacopeia, 1954. Apt. delo 3
no.5:54-57 S-0'54.
(PHARMACOPPIA,
Hungar, 5th edition)

SUNAROKOVA, M.Ta., dotsent.

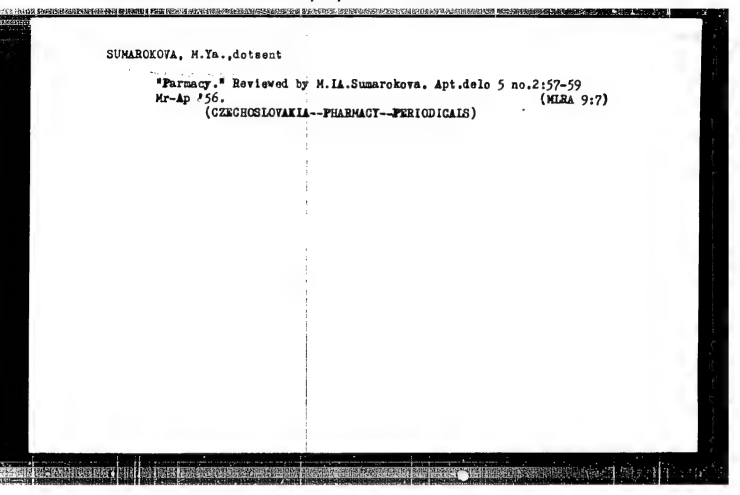
Pharmaceutical education in Csechoelovakia. Apt.delo 4
no.5:59-64 S-0 '55. (MLRA 8:12)
(PHARMACT, education,
in Csech.)

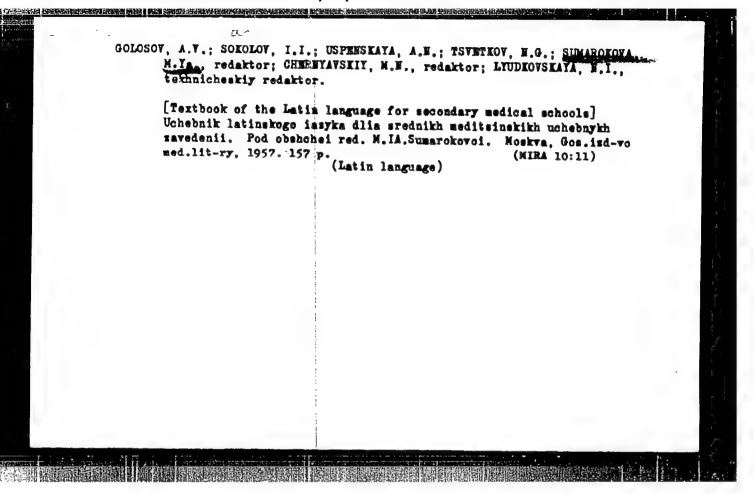
SUMAROKOVA, M.Ya., dotsent.

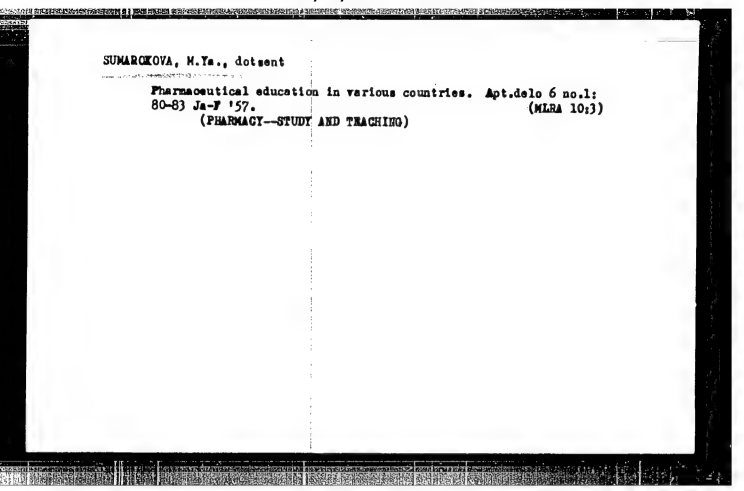
Pirst supplement to the seventh edition of the French Pharmacoposia.

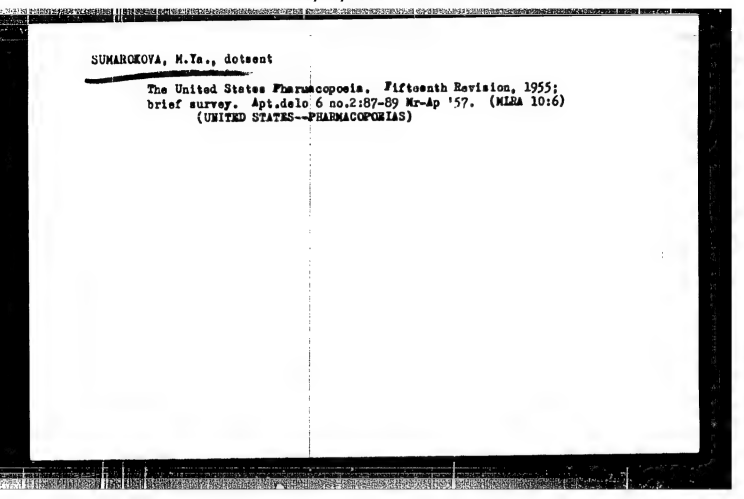
M.Ia. Sumarokova. Apt. delo. 4 no.6:50-52 N-D '55. (MIRA 9:1)

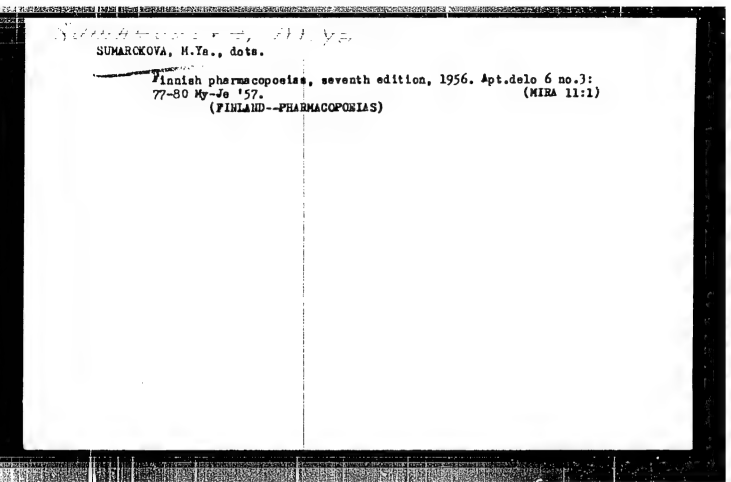
(FRANCE-PHARMACOPOMIAS)

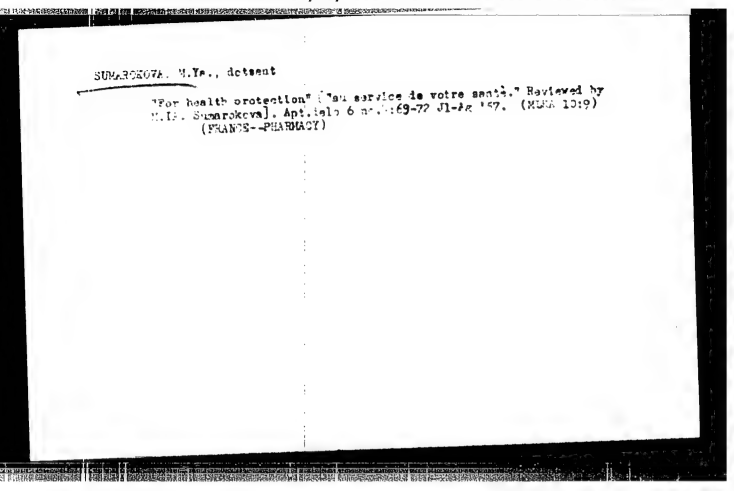


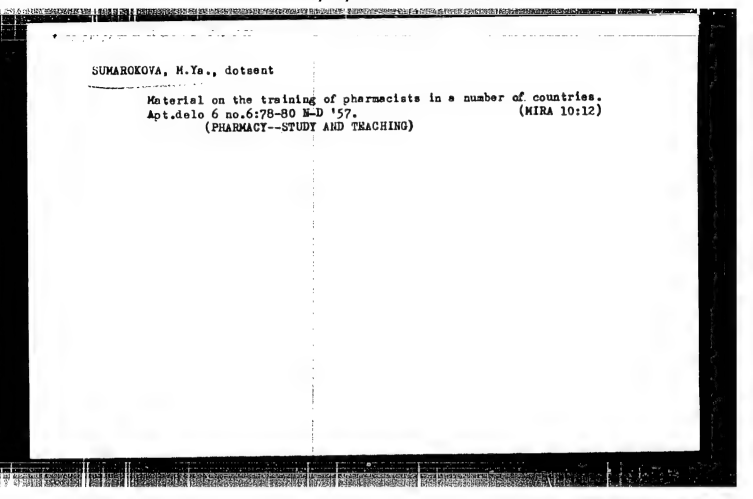


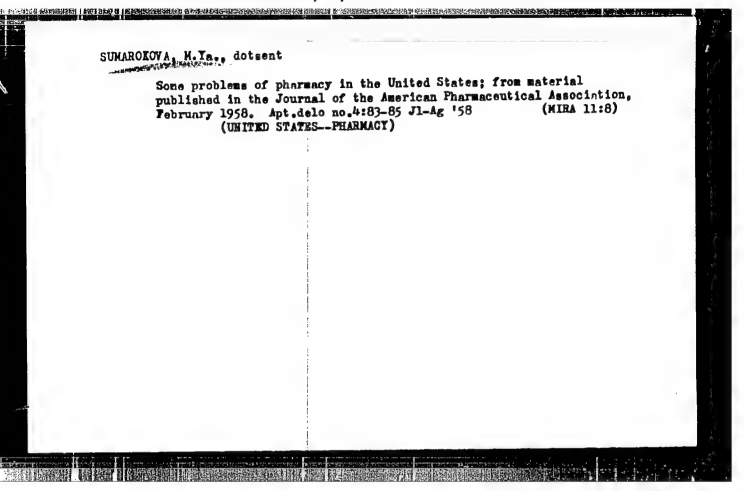


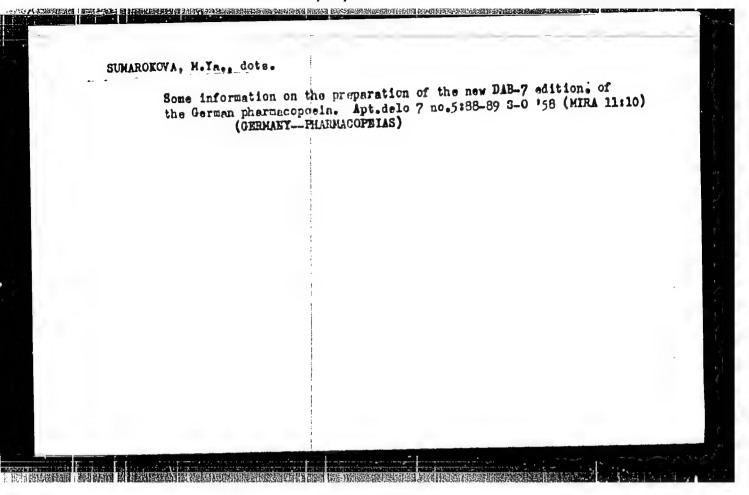


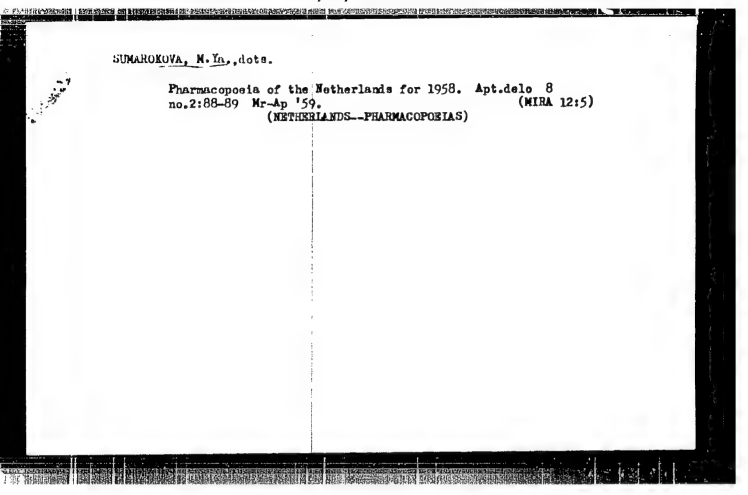












AUTHORS: Kolchin, O.P., Sumarokova, N. V., Chuveleva, N. P., 89-12-5/29

TITLE: Production of Plastic Niobium (Polucheniye plastichnogo niobiya)

PERIODICAL: Atomnaya Energiya, 1957, Vol. 3, Nr 12, pp. 515-524 (USSR)

ABSTRACT: First the properties of niubium are written down. The process necessary for obtaining niobic powder with 90,9 - 99,2% content of niobium is described in detail. This powder is obtained by reduction of the K2NbF7 with sodium. If from the powder obtained pressed bars are manufactured and sintered in the vacuum, plastic niobium is obtained. The investigation of the phase condition of this not entirely reduced mixture of niobic oxide and niobic carbide shows that the main reaction in the reduction in the vacuum can be understood as a summation reaction of successively occurring reactions

in the following form: $Nb_2O_5 + 5 NbC \xrightarrow{\langle 1200 \circ C \rangle} 2 NbO_2 + 5"NbC" (NbCO.8) + CO$

where "NbC" is the phase with variable composition: 2 NbO₂+5"NbC" (NbC_{0,8}) $\sim 1200^{\circ}\text{C}$, 0,5 NbO₂ + 1,5 NbO + 2,5 Nb₂C+1,5 CO 0,5 NbO₂+1,5 NbO+2,5 Nb₂C $\sim 1400^{\circ}\text{C}$, 2NbO+2Nb₂C+Nb+0,5 CO 2 NbO + 2 Nb₂C + Nb $\geq 1400^{\circ}\text{C}$, 7 Nb + 2 CO

Special investigations gave evidence that in the choice of the reduction regimen the interaction between the vapours of the infer-

Card 1/2

S/180/61/000/006/002/020 E021/E135

AUTHORS 5

Sazhin, N.P., Kolchin, O.P., and Sumarokova, N.V.

(Moscow)

TITLE

The processes of reduction of niobium oxides by

carbon

PERIODICAL: Akademiya nauk SSSR. Izvestiya. Otdeleniye tekhnicheskikh nauk. Metallurgiya i toplivo,

no.6, 1961, 8-24

The chemical and physical processes occurring during high temperature reduction of oxides of niobium by carbon were TEXT: studied with the aim of explaining the mechanism of reduction. Niobium pentoxide powder (0.15 mm particles) containing 0.01% TiO2, 0.06% Fe₂0₃ and 0.01% SiO₂ was used with lamp-black or niobiumcarbide as reducing agents. The niobium carbide was prepared by heating a mixture of niobium carbide with lamp black at 1800 °C in it contained 10.2% carbon. The phase a current of hydrogen; composition of the products of incomplete reduction were studied, a thermodynamic analysis of the Nb-O-C system was made and the properties of the oxides of niobium were investigated. Card 1/4

The processes of reduction of ...

S/180/61/000/006/002/020 E021/E135

of diffusion of oxygen and carbon in niobium and the rates of evaporation of the lower oxides of niobium were compared semiquantitatively. From the results of the experiments and from a critical examination of other literature it is shown that the reduction is a multi-stage process, and a mechanism for reduction at temperatures used in practice is proposed. Reduction by niobium carbide at 1100-1300 °C of the pentoxide to the dioxide and partially to the oxide occurs in the main by the generally accepted two-stage scheme with the formation of carbon monoxide. Reduction may also occur at the beginning of the process by interaction in the solid phase between niobium pentoxide and carbon formed by dissociation of the carbide. The second stage of the process is the reduction of the lower oxides of niobium at temperatures above 1500 °C. This takes place in the gaseous phase and depends on the evaporation of the lower oxides which is the reaction limiting the rate of the process. In the last stage of the process, oxygen and carbon diffuse to the surface of the pores in the metal and are evolved as carbon monoxide. The reaction limiting the purification of the metal from dissolved Card 2/4

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The processes of reduction of ... S/180/61/000/006/002/020 E021/E135

oxygen and carbon is the desorption of the carbon monoxide. The reduction of the pentoxide to the dioxide and the dioxide to the oxide can be carried out in a relatively low vacuum or in a current of hydrogen or inert gas. The reduction of the oxide and the removal of the carbon monoxide at temperatures used in practice require a high vacuum. The rate of reduction can be increased by increasing the temperature and rate of carbon monoxide removal. . The maximum temperature possible for each stage is determined by the melting point of the phase most easily The proposed mechanism for the reduction process of niobium can be extended without any radical changes to the reduction process of tantalum and vanadium from their oxides and from mixtures of their oxides and carbides. There are 6 figures, 4 tables and 33 references: 19 Soviet-bloc, 6 Russian translations from non-Soviet publications, and 8 non-Soviet-bloc. The four English language references read as follows:

Card 3/4

The processes of reduction of ... S/180/61/000/006/002/020 E021/E135

Ref.15, G.L. Miller. Tantalum and Niobium, London, 1959, pp 181-187, 283-291.

Ref.21: High Temperature Technology. N.Y.-London, 1956, ed. J.E. Campbell.

Ref.24: F. Holtsberg, A. Reisman, M. Bewry, M. Berkonbilt. The polymorphism of Nb₂O₅. J. Amer. Chem. Soc., 1957, 79, 2039.

Ref.26: R. Orr. High temperature heat contents of tantalum and niobium oxides. J. Amer. Chem. Soc., 1955, 75, 2808-09.

SUBMITTED: March 21, 1961

S/089/61/010/002/012/018 B102/B209

AUTHORS:

Kolchin, O. P., Sumarokova, N. V.

TITLE:

Melting point and other properties of lower niobium oxides

PERIODICAL:

Atomnaya energiya, v. 10, no. 2, 1961, 168-170

TEXT: The present paper is a report on investigations made on NbO and NbO2. NbO2 was produced by reduction of the pentoxide in vacuo at 1300 and 1700°C, NbO by reduction of NbO2. The impurities in the pentoxide (tantalum-silicon-, titanium-, and iron oxides) were removed for the major part during reduction in vacuo. The three obtained niobium dioxide samples still contained 0.02 - 0.04% by weight of C, 0.006% by weight of N and had the composition NbO 1.942,

NbO_{1.956} and NbO_{1.986}. Radiographic structural analysis showed that the product consisted of one phase only with the following lattice parameters: a = 4.82+0.02 kX, c = 2.99+0.02 kX. The three monoxide samples contained 0.04 - 0.06% by weight C and 0.03% by weight N and exhibited the composition NbO_{0.95}, NbO_{1.01}, and NbO_{1.02} Welting point determination: Briquets

Card 1/3

S/089/61/010/002/012/018 B102/B209

Melting point and other ...

(1.5 - 2.0 t/cm2, 10 x 10 mm dross section) were pressed from NbO and NbO powder and electrically heated in containers made of niobium sheet by a graphite heater in vacuo. The melting point of NbO was found to be at 1935°C, that of NbO2 at 2080°C, allowing for an error of ± 15°C. Microand radiographic structural analysis was employed on re-molten NbO for checking the single-phase consistency and to determine the parameter a (4.201 kX). The results concerning NbO2 do not agree with those form Ref. 9 which is said to be due to the fact that in that case NbO2 contained NbO impurities. Determination of volatility and composition of the gaseous phase: Investigations in vacuo (0.5 - 1.10-6 mm Hg) at various temperature showed that both oxides start evaporating at a considerable rate at 1700°C. At 1850°C, all of the monoxide and 45% by weight of the dioxide were evaporated after 4 hours, and after 8 hours also the dioxide was entirely evaporated. It was proved mass-spectroscopically that the gaseous phase above NeO2 contains NbO2 molecules only, i. e. that neither dissociation nor association take place. Investigation of the micro-hardness: Re-molten homogeneous riobium oxide samples had a micro-hardness of

Card 2/3

Melting point and other ...

S/089/61/010/002/012/018 B102/B209

1930 kg/mm², the eutectic in the average one of 794kg/mm² (load of 50 g), the dioxide 1720 kg/mm²; the latter value is inaccurate. Character of electric conductivity: NbO turned out to have metallic conductivity, NbO to be a semiconductor. Oxidation in air: Dioxides (NbO_{2.00} and NbO_{1.92}) produced at 1300 and 1700°C, respectively, and a monoxide (NbO_{0.914}) produced at 1700°C were powdered (grain size 0.15mm) and heated in air at 100 - 300°C; oxygen content was determined. After 6 hours of heating at 100, 150, 200, 225, 250, and 275°C the oxygen content was unchanged in the monoxide and was a little raised in the dioxide at 275°C. However, the surface of the powder particles of both oxides became yellowish already at 150°C and bronze-colored at 200°C. Only after 6 hours at 300°C, both oxides were completely oxidized to the pentoxide. In conclusion, the authors thank L. V. Mel'nikova for having made the metallographic analysis. There are 2 figures and 11 references: 5 Soviet-bloc and 2 non-Soviet-bloc.

SUBMITTED: October 4, 1960

Card 3/3

ACCESSION NR: AP4042350

were sintered at a temperature varying from 1700 to 2100C; compacts 20 x 20 mm were sintered at 1900C or at 2100C in a vacuum of 0.001 mm Hg. In the preparation of alloys, Al, Ti, Zr, V, Ta, No, more complete and W were used as the alloying elements; for removal of carbon during reduction, the oxides were added in an amount 2-5% higher than the stoichiometric. The experiments showed that binary Nb-(4.8-6.2) XHo, Nb-(5.7-24.6) X W, Nb-(4.3-5.8) X V, Nb-2,2% Ti alloys, ternary Nb-(4.25-4.9)% No-(0.87-1.85)% Zr, Nb-(17.0-26.4) X W-(2.07-4.5) X Ti, Nb-5.0X W-2.0X Ta, Nb-(3.0-3.5) X V-(0.4-0.5) X A1 alloys, and quaternary Nb-14.1% W-5.0% Ho-(0.93-1.1)% Zr alloys can be prepared by one or both process Attempts to obtain binary Nb-Al alloys were unsuccessful. quality alloys with a consistently lower content of O, N, and C are obtained by reduction at 2100C. The alloys have a porosity of 40-50%. Subsequent electron beam melting substantially lowers the content of O, N, and C, in some cases without affecting the content of W and Mo, or Zr when its content is about 1%. When necessary, the alloys can be reprocessed by any conventional method used for unalloyed niobium. Experiments on electron beam melting of the alloys were ducted by A. V. Yelyutin. Orig. art. has: 3 tables. Card 2/3

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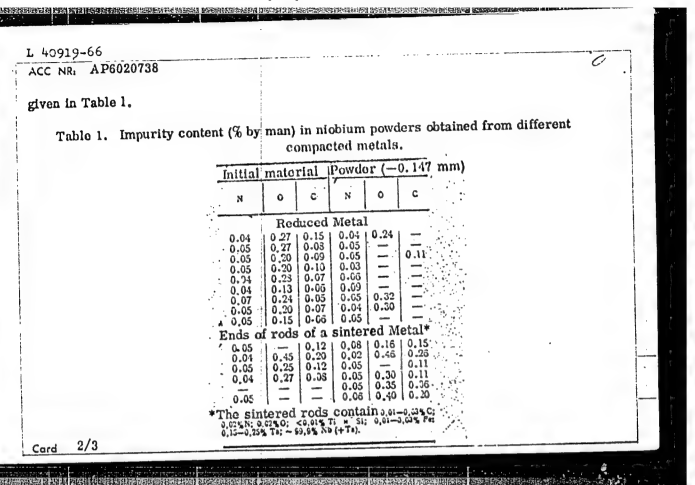
的人,我<mark>是我们的人类的心理,我们都没有的人的人</mark>的人,我们也没有这些的人的人,我们就是这个人的人,我们就是这个人的人的人,我们就是这个人的人,我们们就是这个人的人 JD/JG EWT(m)/T/EWP(t)/ETI [JP(c) L 32686-66 SOURCE CODE: UR/0136/66/000/004/0067/0070 ACC NR: AP6012727 Kolchin, O. P.; Sumarokova, N. V.; Vol'dman, M. A. AUTHOR: ORG: non€ TITLE: Kinetics of the combined carbothermic reduction of niobium and tungsten SOURCE: Tsvetnyye metally, no 4, 1966, pp 67-70 TOPIC TAGS: vacuum furnace, chemical reduction, niobium, tungsten, niobium compound/ /VVPS-10A type vacuum furnace ABSTRACT: This is a continuation of a previous investigation (O. P. Kolchin et al. Tsvetnyye metally, 1964, no 7) with the difference that it deals with a detailed investigation of the kinetics of the combined carbothermic reduction of Nb and W from the mixtures of the oxides and carbides of Nb and alloy elements at various temperatures, the degree of reduction being determined according to the change in the C content of specimens following their heating in a VVPS-10A type vacuum furnace. In the reduction products the W content was determined by the photocolorimetric thiocyanate method, correct to 3-5% (rel.); the Nb content, according to weight gain when heating the specimen in air; and the C content, by the volumetric method. It was found that in the presence of W the degree of the reduction of Nb205+5NbC at 1200 and

Card 1/2

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1400°C i an oxide first to Nb, whic temperat confirm get subl	s greate: rather get red; h is accourse of lathe earl: imated fi	r, partic than a ca uced and ompanied 400°C and ier obser rom the b	rbide. X-ray subsequently, by the reducti regularly inc vation that W	structural at higher on of Nb. rease it u oxides.dur even allo	phase analys temperatures, It is best to ntil at most ing the reduc ys with a high	s 10% W in the sis showed that in it forms an all o start with the 1900°C. These stion virtually on W content have	is the loy with strings lo not
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ACC NR: AP6020738	SOURCE CODE: UR/0136/66/000/006/0065/0067
Men'shchikov, V. A.; Kadyshevskiy	va, N. P.; Sumarokova, N. V.; Filipenko, V. V.; & V. S.; Belimov, N. I.; Abramovich, E. B.
ORG: none	21
TITLE: Manufacture of powdered nic	bium and its alloys by hydrogenating compacted metals
SOURCE: Tsvetnyye metally, no. 6,	1966, 65-67
hydrogenation, niobium alloy	r metal production, niobium, powder metallurgy,
genating niobium or its alloys at low pressures (up to 0.7 atm) than those reduced to appearature levels. Hydrogenetics ource materials derived by electron	method for manufacturing high purity powders by hydro- er temperatures (360 to 400C) and lesser excess hydrogen commonly required. The process is even faster at the genation and milling techniques are given in detail for a beam smelting or carbide heating processes. For the d powder was 91.4%, total yield 98.3%, unaccountable loss- obium powders obtained from different compacted metals is
Card 1/3	UDC: 669.293-492.2



Orig. art. has: 2 figures and 1 table.	!
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. *Incr-61* = wr(n)/1/200(t)/211 -- 1J2(c) -- JD/J6

ACC NR: AP6031728

SOURCE CODE: UR/0136/66/000/009/0072/0074

AUTHOR: Kolchin, O. P.; Filipenko, V. V.; Nizharadze, K. S.; Abramovich, E. B.; Sumarokova, N. V.; Men'shchikov, V. A.

ORG: none

TITLE: Synthesis of niobium carbide with a low nitrogen content

SOURCE: Tsvetnyye metally, no. 9, 1966, 72-74

TOPIC TAGS: niobium carbide, high purity carbide, learnitrogramiohium carbide, niobium carbide synthesis, NICOUM COMPOUND, CAROIDE, NITROGEN, CXYCEN, CHEMICAL SYNTHESIS

ABSTRACT: An investigation has been made of the various factors which contribute to the contamination with <u>nitrogen</u> and <u>oxygen</u> of niobium carbide produced by a continuous process in the Tamman furnace. The investigation results showed that the only significant source of contamination was the inflow of air into the reaction chamber when the furnace was opened every 30 min for charging and removing the final product. Modification of the charge chamber decreased the cross section of the charging shute from 1000 to 160 cm², cut in three the number of openings required to charge the chamber, and sharply reduced the amount of the air flowing in through a narrowed charge shute. A hydraulic lock was also installed for combustion gases, which made it possible to increase the pressure of gases in the furnace to 100—200 mm Hg and thus practically eliminate the inflow of air into the furnace.

Card 1/2

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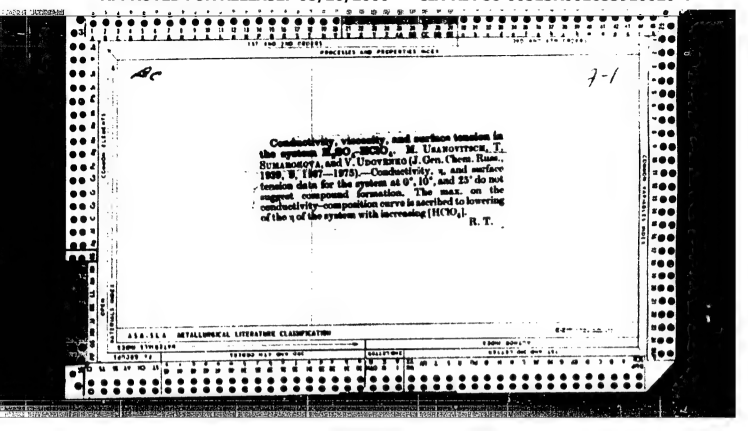
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ACC NR: AP6031728

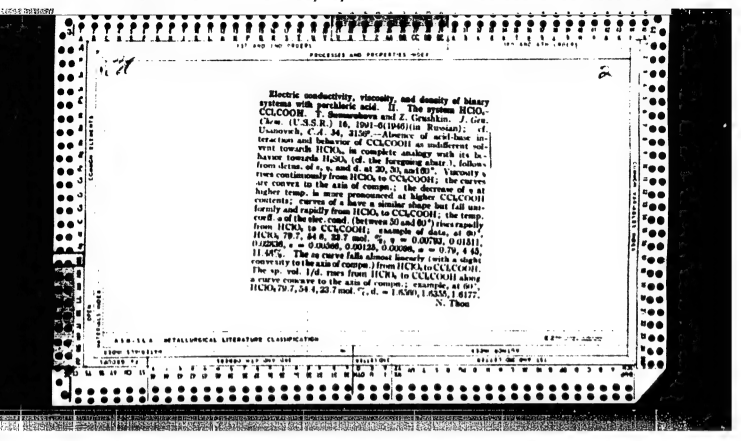
The resulting Improved the quality of niobium carbide produced. The niobium carbide produced in the modernize 53912020-4" contained 89.32—89.63% Nb(+Ta), 0.03—0.14% Fe, 10.0—10.4% C, and only 0.023—0.059% N and 0.14—0.52% O, instead of the previous 0.3% N and 2—3% O. Tantulum carbide with a low content of nitrogen and oxygen was also produced in the modernized furnace, and it is believed that pure carbides of other refractory metals can be produced in it. Orig. art. has: 2 figures.

SUB COUE: 07 / SUBM DATE: none/ ORIG REF: 005/ OTH REF: 001

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SUMARCKOVA, T.

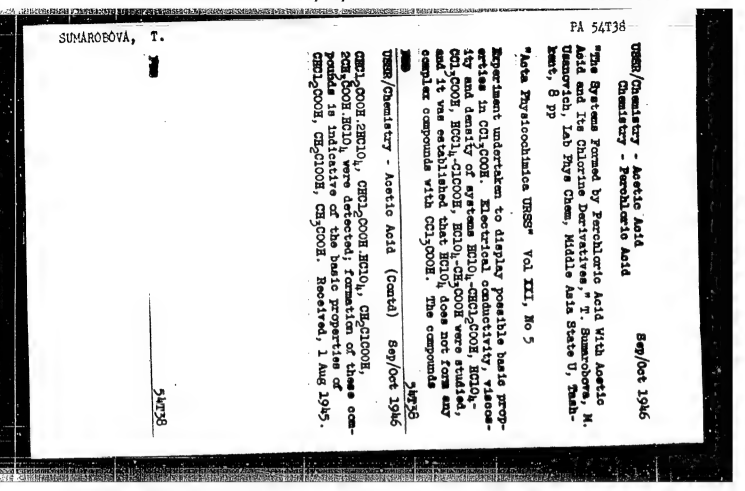
USSE/Electricity Conductivity Conductance - Charts Sep/Oct 46

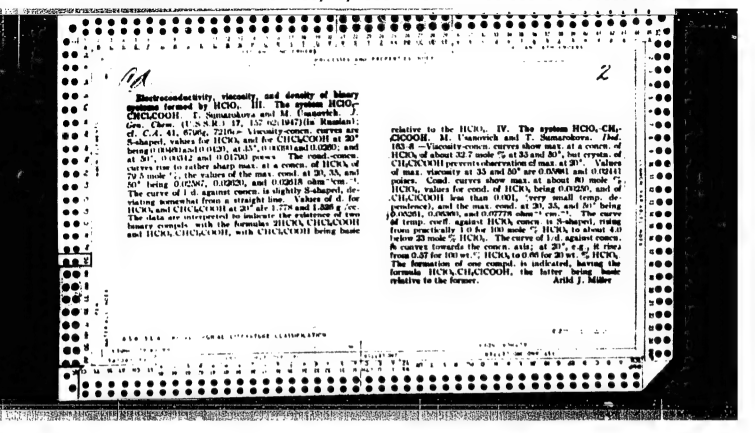
"The Electrical Conductivity of the System HClO, H2O," M. Usanovich, T. Sumarokova, Lab Fhys Chem, Middle Asia State U, Tashkent, 5 pp

"Axta Physicochi:ica URSS" Vol XXI, No 5

Complete electrical conductance diagrams obtained at temperatures 20°, 50°, 60°; and the 50° isotherm investigated at great length. Data obtained indicates chemical interaction in system; hydrates of perchloris acid, HCAO₂cH₂O appear to manifest themselves in liquid phase. Received, 1 Aug 1945.

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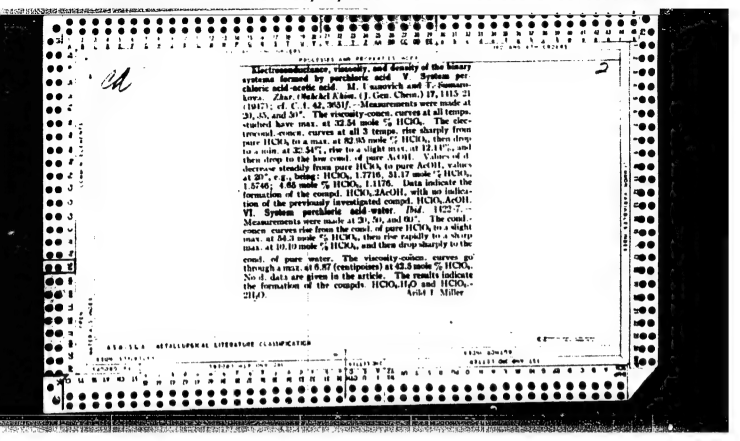




STRICTOWN, T.

Uganovich, M., and Sumarokova, T. -*Electroconductivity, Viscosity and Density of Binary Systems of Binary Systems formed by HClO₄. IV. The System HClO₄.—
CH₂ClOCOH.* (p. 168)

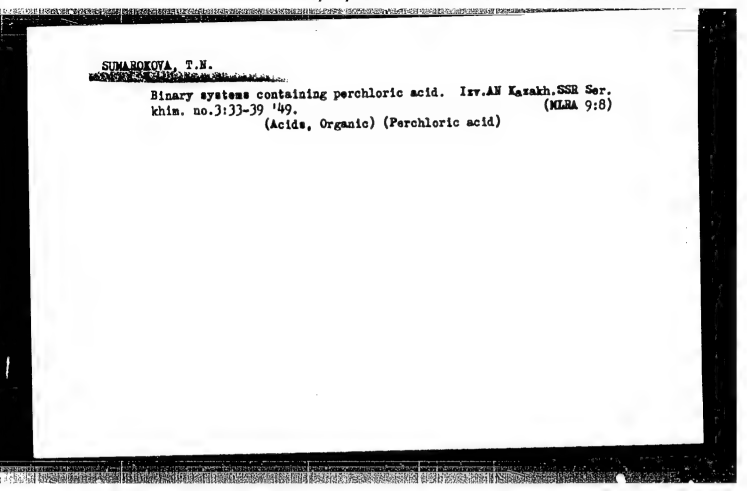
SO: Journal of General Chemistry, (Zhurnal Obshchei Khimii), 1947, Vol. 17. No. 2.



STEAROROVA, T.

Usanovich, M., and Sunarokova, T.- "Electroconductivity, Viscosity and Density of the Binary Systems formed with ECIO4. VI. The System HCIO4-H2O (p. 1427)

SO: Journal of General Chemistry, (Zhurnal Obshchei Khimii), 1947, Vol. 17, No. 8



Strategy, to; Michigan, F.

Actua, Organic

Complex compounds SnCl4, SbCl3, and AsCl3 with some organic acids, Irv. Sekt. plat. i blag. met. Do. 25, 1950.

9. Monthly List of Russian Accessions, Library of Congress, April 1952, Uncl.

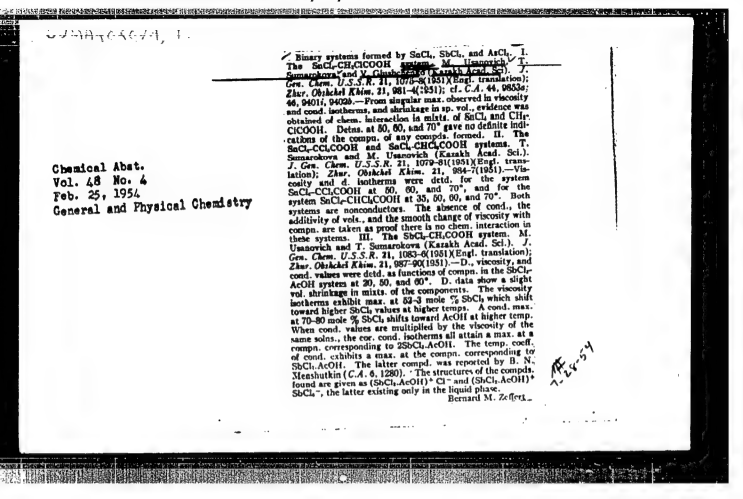
SMALORITA, d.

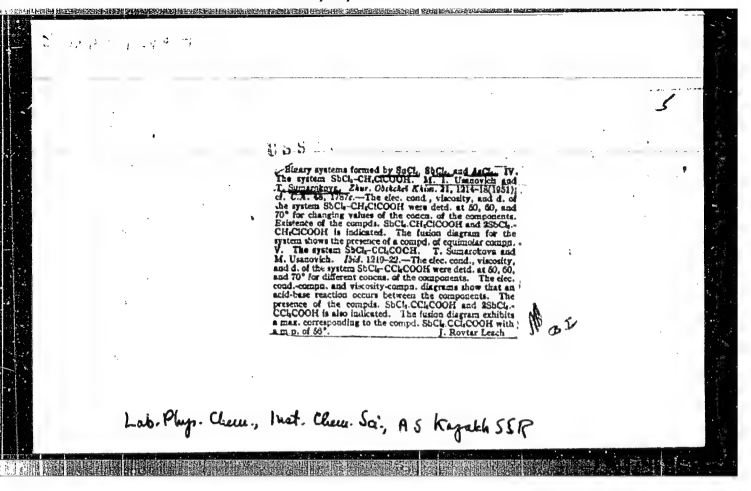
"Rimary systems containing SNOL, Stoll, and Asoll. I. The system Snol,-ch_Clooch." by H. Ucanorich, I. Surrokova, and V. Glushchenko. (p.901)

S0: Journal of General Chemistry (Zhurnal Obshchei Khimii) 1951, Voluen 21, No. 6

SCHEROLIM, 1.

"Binant supplies combaining Smol, Smol, and Asol, II. The system Smol, Coll, Coll

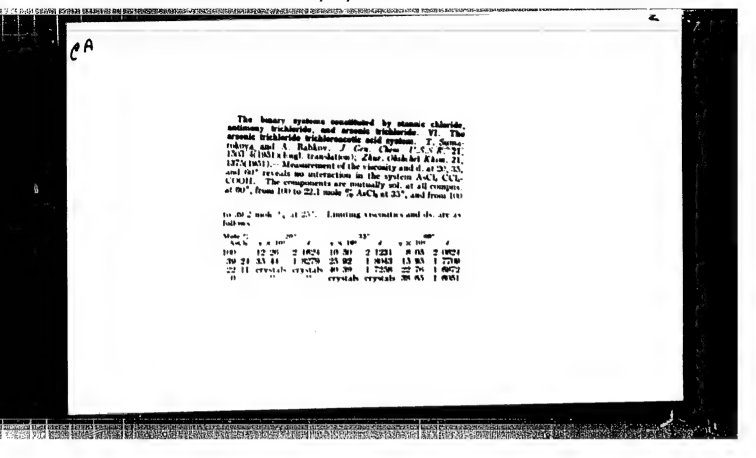




"Binary systems containing SnCl₄. SbCl₃ - AsCl₃. V. The system SbCl₃ - CCl₃CCCH."

T. Sumarkova and M. Usanovich. (p. 1219)

SC: Journal of General Chemistry (Zhurnal Chahchei Khimii) 1951, Vol 21, No 7.



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19:17

JUMILLANOVA, T.

USSR/Chemistry - Arsenic Compounds

Aug 51

"Binary Systems Formed by SnCl₃, SbCl₃, and AsCl₃.
VII. The Systems AsCl₃-CH₂ClCOOH and AsCl₃-CH₂CICOH," T. Sumarolova, V. Glushchenko, Student,
Lab of Phys Chem, Inst of Chem, Acad Sci Kazakh
SSR

"thur Obsheh Khim" Vol XXI, No 8, pp 1376-1380

system AsCl₃-Ch₂ClCCOH, form of isotherms of rescosity and density at temps 50, 60, 70°C and because of elec cond showed that components do not be error. In system AsCl₃-CH₂COOH, study of elected at temps 50,60° and viscosity and density at 20,50,60,70° established that there is acid-base interaction.

SUMAROKOVA, T.N.; LITVYAK, I.G.

Complex compounds SnCl₁₀ · 2A · B and SnCl₁₀ · 2A · 2B. Report no.1.

Ixv.Sekt.plat.i blag.met. no.27:127-136 · 52. (KIRA 7:5)

1. Institut khimicheskikh nauk Akademii nauk Kazakhskoy SSR,
Alma-Ata. (Cempounds, Cemplex) (Tin compounds)

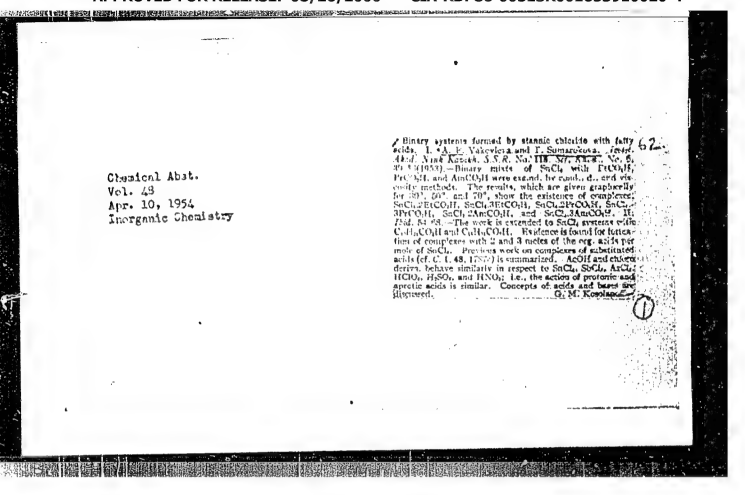
SUMAROKOVA, T.N.; MAKSAY, L.I.

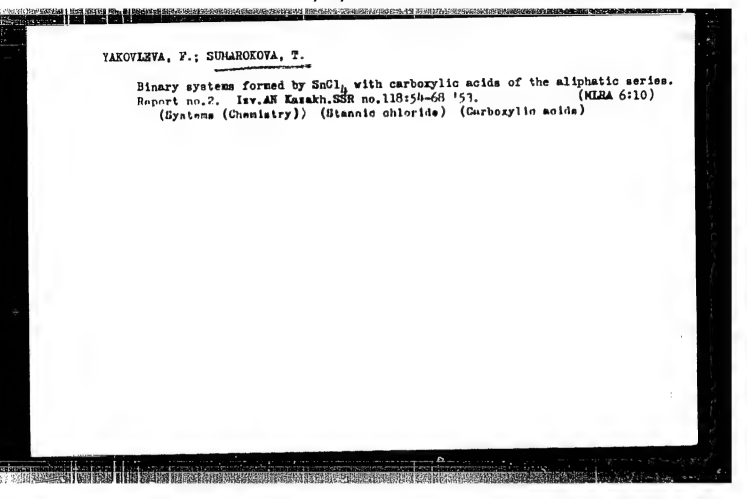
Complex compounds SnCl₁*2A*B and SnCl₁*2A*2B. Report no.2.

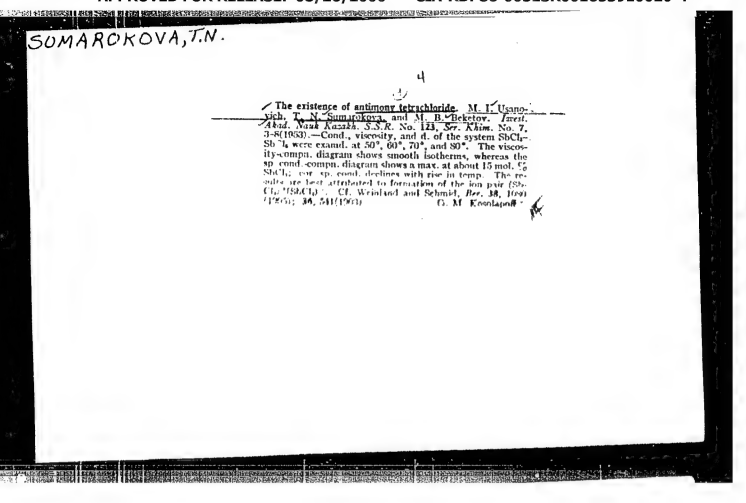
Inv.Sekt.plat.1 blag.met. no.27:137-151 '52. (MERA 7:5)

1. Institut khimicheskikh nauk Akademii Kasakhskoy SSR, Alma-Ata.

(Compounds, Complex) (Tin compounds)



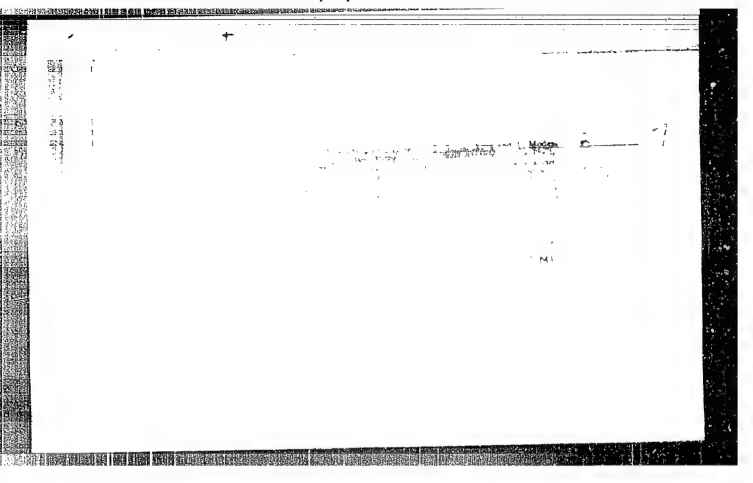


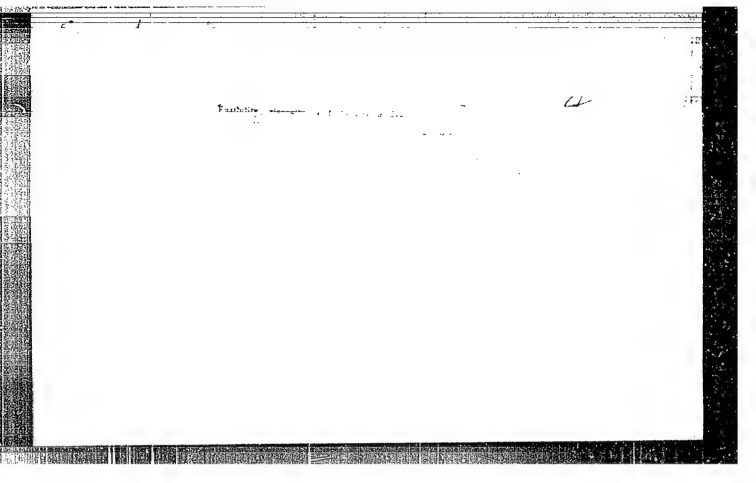


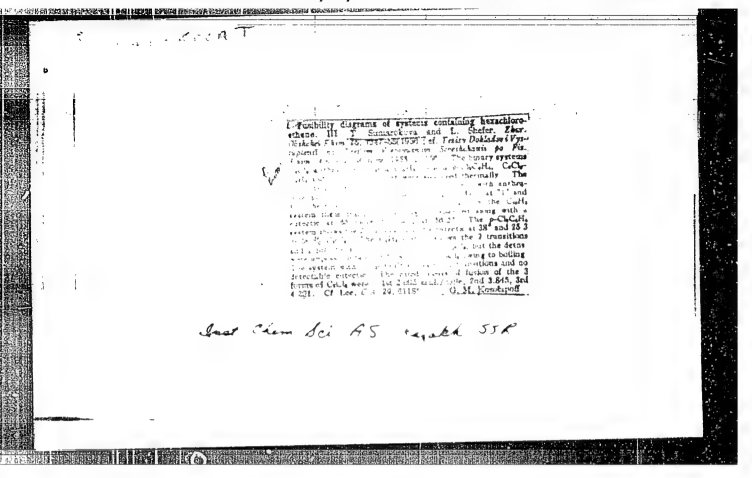
SUMAROKOVA, T.; BOLYALOV, I.

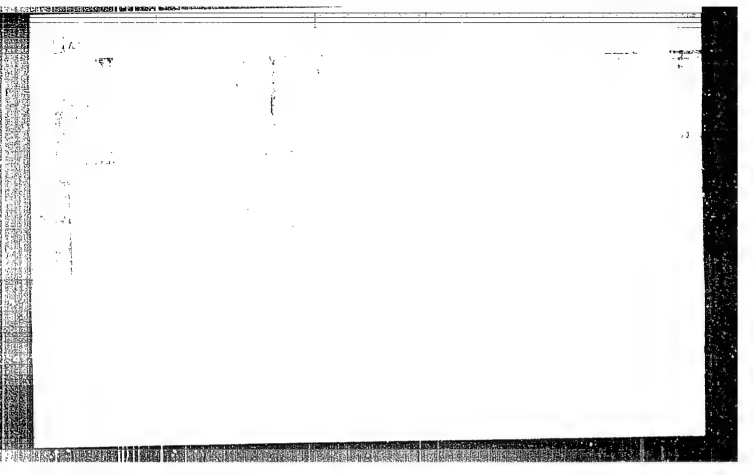
Oxonium compounds of esters with organic acids. Part 2. System: cetyl acetate--acetic acid. Zhur.ob.khim. 25 no.3:477-479 Mr *55 (MLRA 8:6)

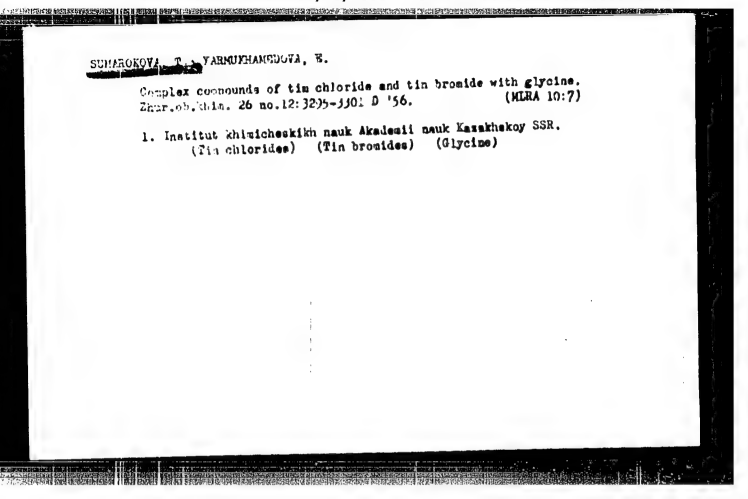
1. Institut khimicheskikh nauk Akademii nauk Kasakhskoy SSR (Acetic acid)(Cetyl acetate)











S UMAROKOVA, T.N.

137-58-2-3907

Translation from: Referativnyy zhurnal, Metallurgiya, 1958, Nr 2, p 233 (USSR)

AUTHOR: 5

Sumarokova, T.N.

TITLE:

Eutectic Diagrams of the Fusibility of Binary Systems. Communication Nr 1 (Ob evtekticheskikh diagrammakh plavkosti

dvoynykh sistem. Soobsheniye 1)

PERIODICAL: Izv. AN KazSSR. Ser. khim., 1957, Nr 1, pp 3-11

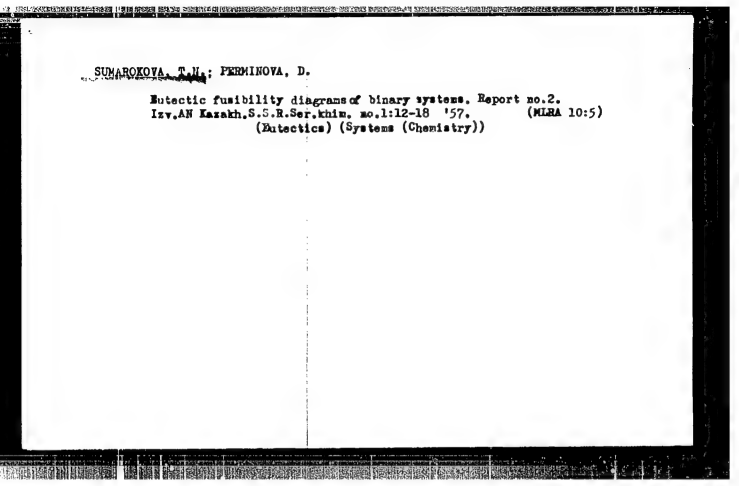
ABSTRACT:

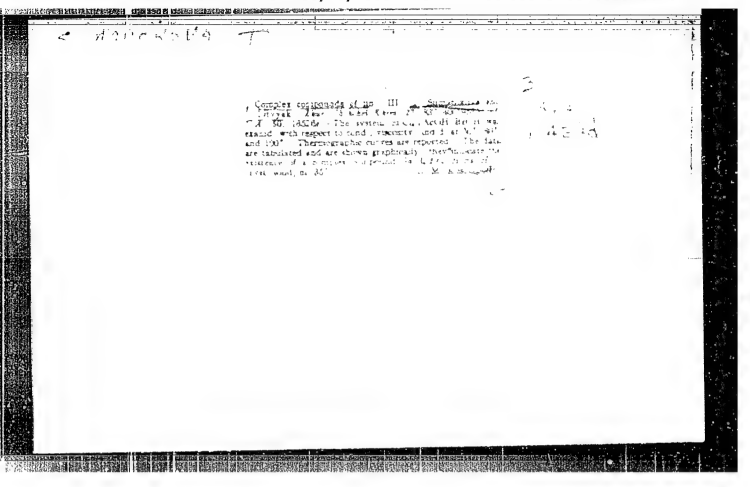
A check was made of the applicability of the Schroeder equation to real systems (organic substances, metals, salts) on the basis of available experimental data from the literature. It is shown that the Schroeder equation is valid at the most varied concentrations in different systems, and that many real systems are subject to the laws of ideal solutions. In ideal binary systems containing a common component, the liquidus curve is the geometric locus of the eutectic points. It is shown that molecular weight may be determined from the data of thermal analysis and the Schroeder equation.

A.F.

Card 1/1

1. Molecular weight—Determination 2. Schroeder equation—Applications





SUMAROKO7A, T.; LITVYAK, I.

Complex tin compounds. Part b. Zhur.ob.khim. 27 no.5:1125-1130
Zhur.ob.khim. 27 no.5:1125-1130 My '57. (MLRA 10:8)

1. Institut khimii Akademii nauk Kasakhskoy SSR.

(Complex compounds)

(Tin organic compounds)

"APPROVED FOR RELEASE: 08/26/2000 CIA-RDP86-00513R001653910020-4

79-12-41/43 Sumarokova, T., Nevskaya, Yu. AUT HORS: Complex Compounds of SnCl₄, Sn Br₄ and TiCl₄ With Cinecle (Kompleksnyye soyedineniya Sn Cl₄, Sn Br₄ i MILE: TiCl_A s Tsineolom). Zhurnal Obshchey Khimii, 1957, Vol. 27, Nr 12, PURIODICAL: pp. 3375-3379 (USSE) The complex compounds SnCl4, Sn Br4 and TiCl4 with organic ABSTRACT: oxides have been little investigated. Compounds of SnCl and SnBr4 with dioxane, as well as of SnCl4 with lactones $(SnX^2 2 A)$ are described in publications. The authors were interested in the systematical investigation of the complex compounds Sn+4 and T+4 with such organic compounds, which possess in their molecule the group -COC-. They chose cincole, which is a constituent part of many etheric oils. The complex compounds of the tin- and titanium halides with cineole were of an extended interest, in as much they are connected with the research to find a method for the quantitative determination of cineole. Mixtures of SnCl4, SnBr and TiCl4 in an exact molecular ratio with Card 1/3

Complete Compounds of SnCl₄, SnBr₄ and FiCl₄ With Cineole 79-12-41/43

cincole were prepared for the synthesis of these compounds. In the course of the reactions a considerable amount of heat was liberated. The components were mixed in indifferent solvents. The composition of the complex compounds was determined analytically and according to the crysoopic method of titration. The amount of tin and titanium was computed as SnO2 and PiO2 the halides were determined according to Vollhard and the Cryoscopic measurements were conducted according to Beckmann. The authors put up diagrams on the basis of the results, from which the dependence of the depression (or of the molecular weight) on the composition, expressed in molecular percent, may be read. The following complex compounds were synthesized: $SnCl_4$. $2C_{10}H_{18}C$, SnB_4 . $2C_{10}H_{18}O$ and TiCl $_4$. $2C_{10}^H{}_{18}$ O. The compound $SnBr_4$. $2C_{10}^H{}_{18}^O$ possesses quite extraordinary properties: It shows a molecular weight of 746'8 and destillates already at 35°C. There are 3 figures, and 7 references, 5 of which are Slavic.

Card 2/3

"APPROVED FOR RELEASE: 08/26/2000 CIA-RDP86-00513R001653910020-4

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Complex Compounds of SnCl₄, SnBr₄ and TiCl₄ Jith Cincole 79-12-41/43

ASSOCIATION: Institute of Chemistry AS Kazakh SSR (Institut khimii Akademii nauk Kazakhskoy SSR).

SUBMITTED: October 31, 1956

AVAILABLE: Library of Congress

1. Complex compounds - Synthesis

Card 3/3

On the Electrolytic Dissociation of Tin and Antimony Complex Compounds.



acid and ethyl acetate, where the organic compound contained the isotope C14. In each case SnCl4 was carried to the anode as well as to the cathode. This harmonizes with the equations 1 and 2. While this paper was being written, two essays were published (Miskidzh'van; Kuz'mina and Vol'nov) in which the authors suggest own schemes of the electrolytic dissociation of complex compounds. These schemes have in common that the formation of complex compounds is meant to represent an incorporating reaction, i.e. the molecule of the organic matter is to be a component of the complex cation. The experiments of the authors of the present paper contradict these schemes. For only the complex compounds of the 3-halides of antimony (and arsenic) with organic oxigen containing substance are incorporation products. The conpounds of the same organic substances with the 4-halides of tin are not incorporation products. In the case of electrolysis according to the equation 3 the authors of the present paper have added methyl-red in addition to the acid marked with C14. As expected, the pigment moved into the same direction as the acetic acid, that is towards the cathode. (With 8 citations from publications).

Card 2/3

"APPROVED FOR RELEASE: 08/26/2000 CIA-RDP86-00513R001653910020-4

POLOSUKHIN, Porfiriy Porfir'yevich, saslushennyy master sporta. Prinimal uchastiye: REVZIN, Sergey Vladinirovich, inzh.-vozdukhoplavatel. SUMAROKOVA, T.N., red.; MANINA, M.P., tekhn.red.

经代表的情况处理的 建物溶液 用血 计多数形式 医皮肤 计设置 经公司 经国际股份 经国际股份 计

[Notes of an anateur navigator and parachutist; as told to Sergei Revzin] Zapiski sportsmena-vozdukhoplavatelis i parashiutista. Literaturnaia zapis! Sergeia Revzina. Izd.3.. dop. i perer. Moskva, Gos.izd-vo "Fizkul!tura i sport," 1958. 230 p.

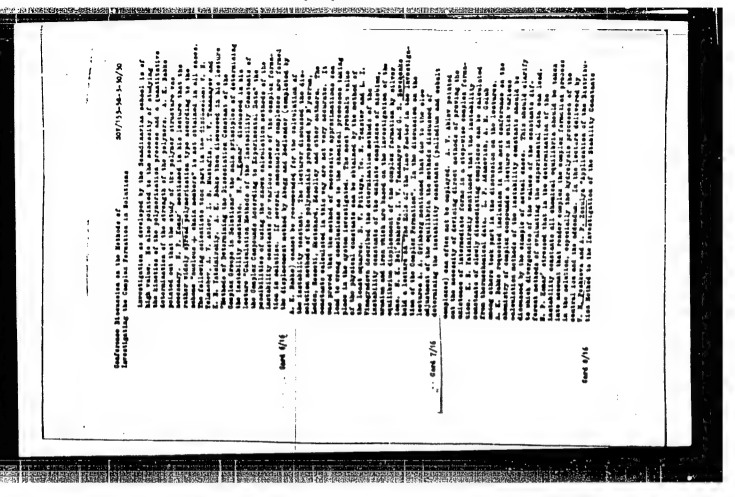
(MIRA 12:12)

(Polosukhin, Porfirii Porfirievich, 1910-)

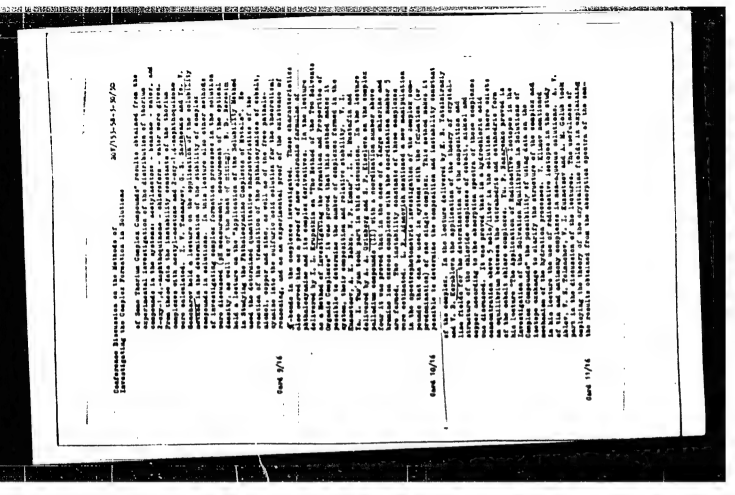
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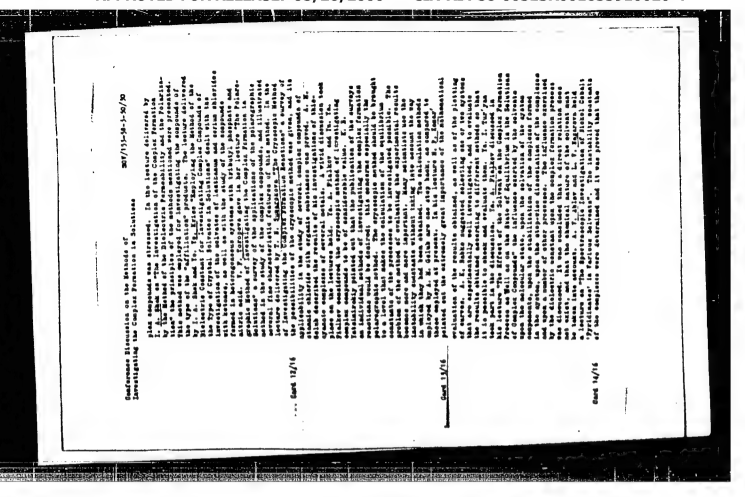
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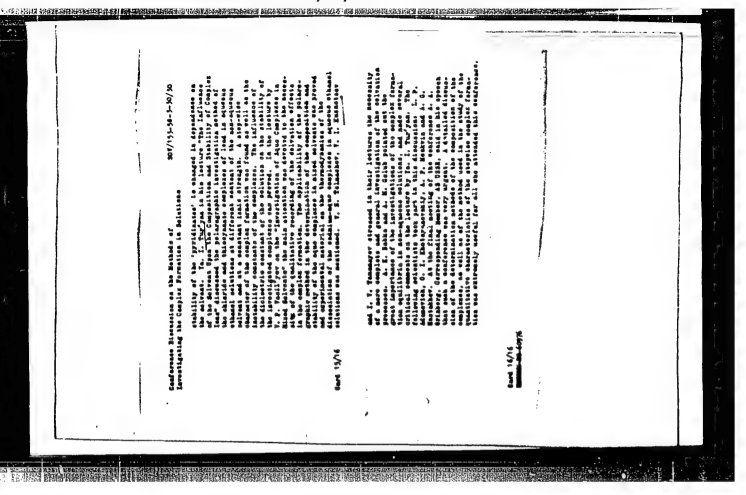


CIA-RDP86-00513R001653910020-4





CIA-RDP86-00513R001653910020-4



307/ 78-3-7-33/44 Možeshova, T., Sumatokova, T. AUTHORS: The Systems PbCh2 PbBz, and PbBz2 PbI2 (Sistemy PbCl2 PbBz2 i TITLE: PhBrg PhJg) Zhastal machganicheakay khimii 1958, Vol. 3, Nr 7, pp. 1655-1660 PERIOD ...AL (USSR) The ayetems PoCl₂ PoBn₂ and PoBn₂ PoT₂ were investigated by means ABSTRACT: of thermal analysis. The compound PhBro was found to occur in form of two modifications. At 3440 C a prose onengof PtCl2 480° C. PbBr2 = 364° C and Pol2 = 396° were determined. The solid solutions of PbCl2 and PbBr2 indicate the existence of the equimolar compound PtCBr. The phase transformation of liquid solutions into solid of emodition tion and the transformation of solid solutions of the of modification into solid solutions of the filmodification characterize the melting diagram of the systems. In the system PhiBro Pril there also saist & and B phase are Them are 2 figures: 2 hables and it references, 6 of 28 32 8 3/1 ExaD which are Sommet.

"APPROVED FOR RELEASE: 08/26/2000 CIA-RDP86-00513R001653910020-4

The Systems PbCic PbBrc and PbBrc Pb'c SUV/78-3-7-33/44

ASSOCIATION: Institute Animal An Karakhakay SSR (Institute of Chemistry, AS Karakhakaya SSR)

SUBMITTED: Jake 17. 901

1 Lend bronide-land incide systems -- Analysis 2 Lend bromide - land chiloride systems -- Analysis

Card 2/2

Yarmukhamedova, E. Sh., Sumarokova, T. N. 79-28-5-67/69

AUTHORS:

TITLE:

Complex Compounds of Tin Chloride and Tin Bromide With Urea

(Komplekanyye sojedinenija khlornogo i bromnogo olova s

mochevinoy)

PERIODICAL:

Zhurnal Obshchey Khimii, 1958, Vol 28, Nr 5,

pp 1410 - 1412 (USSR)

ABSTRACT:

In the systematic investigation of complex compounds of halides of tetravalent tin with organic compounds containing nitrogen and oxygen, the authors found that thioures enters reaction with tin chloride and tin bromide with formation of a complex compound of the composition SnX4.2(NH2)2CS (Reference 1). It

was of interest for the authors to experience by investigations which way urea would react on the halides of tetravalent tin. Taking into adcount the similarity of urea with thiourea it was assumed that also the had to form compounds of the same kind with tin chloride and tin bromide. Tin chloride was purified by repeated distillation and the fraction with the boiling temperature 109°C (690°mm) was stored in sealed ampoules. Tin bromide was treated the same way. The complex com-

Cord 1/2

79-28-5-67/69

Complex Compounds of Tin Chloride and Tin Bromide With Urea

pound of tin chloride with urea was obtained by direct action of tin chloride on it without solvent - that of tin bromide the same way. Thus the complex compounds of tin chloride and tin bromide with urea of the following composition were synthetized: SnCl₄·2(NH₂)₂CO and SnBr₂·2(NH₂)₂CO. These complex compounds

are crystalline products, do not change in air or dissolve easily in organic media. There are 2 figures and 1 reference, 1 of which is Soviet.

ASSOCIATION: Institut khimicheskikh nauk Akademii nauk Kazakhskoy SSR

(Institute for Chemical Sciences, AS Kozakh SSR)

SUBMITTED: April 19, 1957

Op. 20 2/2

"APPROVED FOR RELEASE: 08/26/2000 CIA-RDP86-00513R001653910020-4

AUTHORS:

Sumarokova, T. H., Arsenov, G. I.

76-32-5-32/47

TITLE:

Methods of Cryoscopic Measurements (K metodike krioskopiches-

kikh izmereniy)

PERIODICAL:

Zhurnal fizicheskoy khimii, 1958, Vol. 32, Nr 5, pp.1153-1154

(USSR)

ABSTRACT:

The usual method of determination based on the measurement of the maximum temperature rise gives wrong results, as can be seen from a given diagram, as the measuring values do not correspond to the real freezing point of the solutions. In determinations according to M. Bakeyev (Ref 2) which are based on the determination of the heating curves the error by undercooling is removed, however, errors due to insufficient mixing occur. An electromagnetic arrangement of mixing is described, using a multivibrator with two lamps L,-62h8 and L2-6 P3 as well as a transformer EIS-2. In order to improve the distribution of the crystals the cryoscope has an elevated bottom. In order to determine the freezing temperature the heating curves are taken by plotting the temperature on the ordinate and the time on the abscissa. By

Card 1/2

76-32-5-32/47

Methods of Cryoscopic Measurements

means of an example, a solution of benzylalcohol in benzene, the obtained diagrams are represented, the melting temperatures (crystallization) being determined by the point of intersection of two straights. The described electromagnetic mixer can also be used for other purposes. There are 4 figures and 3 references, 3 of which are Soviet.

ASSOCIATION: Akademiya nauk Kazakhskoy SSR, Institut khimii, Insitut energe-

(Institutes of Chemistry and Power Engineering, AS Kazakhskaya SSR)

May 3, 1957 SUBMITTED:

2. Freezing points--Determination 1. Liquids--Freezing

Card 2/2

CIA-RDP86-00513R001653910020-4" APPROVED FOR RELEASE: 08/26/2000

QUKOV, Valentin Ivanovich; SUMAROKOVA, T.B., red.; FEKLISOVA, T.D., tekhn.red.

[In the land of untsuched treasures] V krain netrematykh sekrovishch. Maskva, Gos.isd-ve "Fiskul'tura i spert."

1959. 86 p.

(Altai Termitery--Description and travel)

sov/79-29-5-5/75

5(4) AUTHORS: Sumarokova, T., Omarova, R.

On the Interaction of Tin Chloride With Esters. 1. (O vzaimodeyst-

TITLE:

vii khlornogo olova so slozhnymi efirami.1.)

PERIODICAL:

Zhurnal obshchey khimii, 1959, Vol 29, Nr 5,

pp 1430 - 1437 (USSR)

ABSTRACT:

In the present paper the systems SnCl₄ - CCl₃COOC₂H₅ and Snc14 - CH2C1C00C2H5 by means of physico-chemical analysis conductiometry, viscosimetry, volumetry and cryoscopy were investigated. The operational methods, the preparation and purifica-

tion of tin chloride were described previously (Refs 12 and 16). The results obtained by determination of the viscosity and the

values B galculated from the equation

(Refs 17-19) are presented in table 1. Table 2 gives the determination results of the density. In figure 1 the diagrams property - composition of the system SnCl₄ - CCl₃COOC₂H₅ are

shown. The determination results of viscosity, specific conducti-

Card 1/3

On the Interaction of Tin Chloride With Esters.1.

sov/79-29-5-5/75

vity and density for the system SnCl4 - CH2ClCOOC2H5 are given in table 3 and 4, the calculated values of the corrected conductivity, temperature coefficient of the conductivity, and of the

constant B are presented in table 5. Figure 2 gives the diagrams property - composition of this system. On investigating the behavior of tin chloride with respect to ethyl trichloro- and ethyl monochloro-acetate the following was found: In the system SnCl₄ - CCl₃COOC₂H₅ the components react with one another until the compound SnCl₄.2CCl₃COOC₂H₅ is formed. In this system the

electric conductivity is practically not existing. In the system SnCl₄ - CH₂ClCOOC₂H₅ the reaction between the components is carried on up to the formation of the compounds

SnCl4.2CH2CClCOOC2H5 and SnCl4.3CH2ClCOOC2H5. The latter determines the electric conductivity of the system. As can be seen from figures 4 and 5 the complex acids SnCl4.2RCOOH are con-

siderably stronger than SnCl₄.2RCOOR'. From the comparison of

Card 2/3

"APPROVED FOR RELEASE: 08/26/2000 CIA-RDP86-00513R001653910020-4

On the Interaction of Tin Chloride With Esters. 1.

SOY/79-29-5-5/75

the behavior of ethyl- and ethyl-monochloro-acetate with the behavior of ethyl-trichloro-acetate with respect to tin chloride it can be seen that the electric conductivity then appears in the systems formed from tin chloride and esters, if the compound SnCl_A. 3RCOOR' is formed in the solution. There are 5 figures, 5 tables, and 24 references, 19 of which are Soviet.

ASSOCIATION: Institut khimicheskikh nauk Akademii nauk Kazakhskoy SSR (Institute of Chemical Sciences of the Academy of Sciences, Kazakhskaya SSR)

SUBMITTED:

April 18, 1958

Card 3/3

在自己的时代的 地名美国西班牙克斯 化多元分类 网络拉拉斯 异双甲甲基苯甲基苯甲基甲基甲基甲基甲基

sov/79-29-5-6/75

5(4) AUTHORS: Sumarokova, T., Omarova, R., Kuzimenko, N.

TITLE:

On the Interaction of Tin Chloride With Esters. 2. (0 vzaimodeystvii khlornogo olova so slozhnymi efirami. 2.)

PERIODICAL:

Zhurnal obshchey khimii, 1959, Vol 29, Nr 5, pp 1437 - 1442 (USSR)

ABSTEACT:

In the present paper the influence exercised by the length of the aliphatic alcohol radical upon the acid-basic properties of esters was investigated. Viscosity, density and electric conductivity of the systems SnCl₄ -CH₃COOC₈H₁₇ and SnCl₄ - CH₃COOC₁₆H₃₃

were investigated. The results obtained on the determination of the properties as well as the calculated temperature coefficients of the electric conductivity, the corrected conductivity and the constant B for the system SnCl₄ - CH₃COOC₈H₁₇ which was in-

vestigated at 25 and 50°, are given in table 1. In figure 1 the diagrams property - composition are presented. By physico-chemical analysis it could be concluded that the components of the system react with one another, thus forming a complex compound $SnCl_4.2CH_3COOC_8H_{17}$. The system $SnCl_4$ — $CH_3COOC_16H_{33}$ was in-

Card 1/3

nganataran markaranga kanambanah kanama makatan

On the Interaction of Tin Chloride With Esters. 2.

SOV/79-29-5-6/75

vestigated at 40, 50, 60 and 70°. The determination results are listed in tables 2 and 3, the calculated values of the corrected electric conductivity, the temperature coefficient of the conductivity and the constant B in table 4. The diagrams property composition can be seen in figure 2. On the strength of the physico-chemical analysis it could be concluded that a complex compound SnCl4.2CH3COOC16H33 is formed in the system. It was separated in free state. Its melting point is 56°. The electric conductivity in the system indicates the existence of this compound. Figure 3 compares the values of the electric conductivity at 50° in the systems SnCl₄ - CH₃COOC₂H₅ (Ref 4), $\operatorname{SnCl}_4 - \operatorname{CH}_3 \operatorname{CCOC}_8 \operatorname{H}_{17}$ and $\operatorname{SnCl}_4 - \operatorname{CH}_3 \operatorname{COOC}_{16} \operatorname{H}_{33}$. The value of the corrected conductivity of cetyl acetate solutions is seen to be far smaller than in octyl acetate and especially ethyl acetate solutions. This can be explained by the fact that the formation of the complex compounds SnCl 4.3RCOOR is a secondary redox reaction. It proceeds via the stage of the formation of complex acids SnCl . 2RCOOR which become weaker on lengthening

Card 2/3

"APPROVED FOR RELEASE: 08/26/2000 CIA-RDP86-00513R001653910020-4

On the Interaction of 'Tin Chloride With Esters. 2.

是一种的人,但是一种的人,我们也是一种的人,我们也不是一种的人,我们就是一种的人,我们就是一种的人,我们就是一个人,我们就是一个人,我们可以是一个人,我们可以不

SOV/79-29-5-6/75

of the radical. A similar rule was found in the systems formed from tin chloride and carboxylic acids (Refs 16,17). There are 3 figures, 4 tables, and 17 references, 13 of which are Soviet.

ASSOCIATION: Institut khimicheskikh nauk Akademii nauk Kazakhskoy SSR

(Institute of Chemical Sciences of the Academy of Sciences,

Kazakhskaya SSR)

April 18, 1958 SUBMITTED:

Card 3/3

sov/76-33-1-31/45 Burarokovo, T. H., Fialkov, Ya. A., 5(4) (Alma-Ata), Deceased AU MIONS: On the Cryoscopic Method of the Physico-Chemical Analysis (Classification of Diagrams) (O kripskopicheskom metode fiziko-khimicheskogo analiza (klassifikatsiya diagramm)) TITLL: Zhurnal fizicheskoy khimii, 1959, Vol 33, Nr 1, pp 184-191 PERIODICAL: (USSR) The first method of this type has been worked out by V. V. Udovenko (Refs 1-7) and is based on the establishment of the function between the molecular weight of the components ABSTRACT: (and their mixtures) of the concentration in the cryoscopic solvent and the marking of the intersection points of the isoconcentration in the coordinates molecular weightcomposition. For plotting the diagram composition-properties by the cryoscopic method of the physico-chemical analysis of the system A-B-solvent, N. A. Izmaylov (Refs 6-11)(and others (Refs 12, 13)) suggested the establishment of the deviation from the additive depression on the ordinate of properties. Ya. A. Fialkov and I. D. Muzyka (Ref 16) used the depression change as a property and based the determination on a Card 1/3

On the Cryoscopic Method of the SOV/76-33-1-31/45 Physico-Chemical Analysis (Classification of Diagrams)

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measurement of the depression of the solutions which corresponds to the intersection point of a pencil of lines in the diagram depression change-composition (Fig 1). The use of precipitation reactions in cryoscopic investigations brought about a considerable increase in analytic possibilities (Ref 17). The amount of the depression is connected to the concentration by the equation according to Schröder (Shreder) (1). On account of (1) the diagram of the ideal case is established and the diagram types depression-composition of systems with a chemical reaction are investigated. It is stated that (in this case) there are three basic types of diagrams. The first type shows a decreasing depression value, the second type a constant value, and the third type a value increasing up to the end point of the depression. A number of examples is given illustrating the types mentioned

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sov/76-33-1-31/45 Physico-Chemical Analysis (Classification of Diagrams) On the Cryoscopic Method of the

(e.g. $SnCl_4 = C_6 + A_2 = C_4 + C_2 = C_4 + C$ in benzene (Hef 19), $\operatorname{lin}(C_2 I_5)_2 - C_3 II_5 NCS$ in benzene (Ref 16),

NH₂CH₂COOH - CCl₃COOH in accetic acid, etc). Several

deficiencies of a previous paper (Ref 8) are pointed out.
There are 3 figures and 21 Soviet references.

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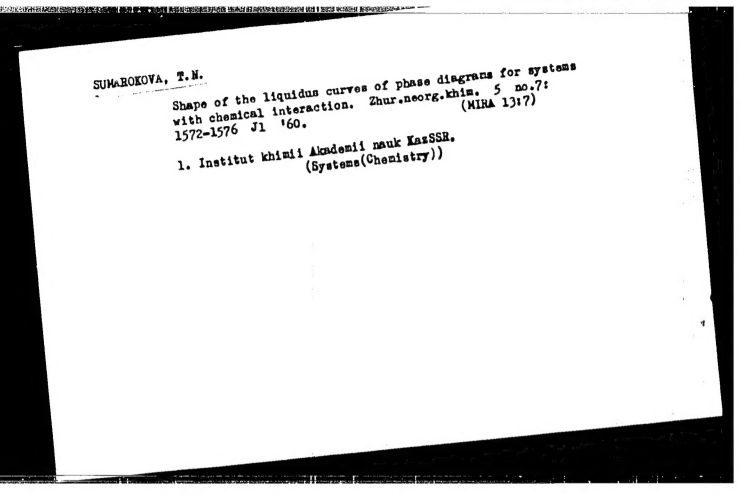
Academy of Sciences bkrSdd (natitute of seneral and inorganic

Chemistry)

July 8, 1957 SUBMITTED:

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"APPROVED FOR RELEASE: 08/26/2000 CIA-RDP86-00513R001653910020-4



SUMAROKOVA, T.N.; MODISTOVA, T.P.

Fusibility in the system Pb61 - KCl. Zhur. neorg. khim. 5 no.11: 2477-2482 N '60. (MIRA 13:11)

1. Institut khimii Akademii nauk Kazakhskoy SSR.
(Lead chloride) (Potassium chloride)